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Drivers and mechanisms of tree mortality in moist tropical forests

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I) Summary Tree mortality rates appear to be increasing in moist tropical forests (MTFs) and could have significant consequences for the global carbon cycle. We review the state of knowledge regarding the drivers of MTF tree mortality, create a conceptual framework with testable hypotheses regarding the drivers, mechanisms, and interactions that may underlie increasing MTF mortality rates, and identify next steps for improved understanding and reduced predictive uncertainty. Increasing mortality rates are associated with rising temperature and vapor pressure deficit, liana abundance, drought frequency, wind events, fire, and possibly CO₂ fertilization-induced increases in stand thinning or acceleration of trees reaching larger, more vulnerable heights. These mortality drivers alter plant physical structure or physiological processes such as carbon starvation and hydraulic failure. The relative importance of each driver is unknown and collectively they are a major knowledge gap. High species diversity may buffer MTFs against large-scale mortality events relative to the extratropics, but the historical and expected trends in mortality drivers give reason for concern regarding increasing mortality within MTFs. Models of tropical tree mortality are advancing representation of hydraulics, carbon, and demography, but require more empirical knowledge regarding the most common drivers and their subsequent mechanisms. We outline critical datasets and model developments required to test hypotheses regarding the underlying causes of increasing MTF mortality rates, and improve prediction of future mortality under climate change.

II) Introduction

Moist tropical forests (MTFs, see Glossary) are the largest terrestrial carbon sink in the world (Pan et al. 2011) and house the majority of Earth's terrestrial biodiversity (Myers et al. 2000, Kreft & Jetz 2007). The spatial patterns of biomass carbon storage in MTFs are primarily driven by mortality (see Glossary) rather than productivity (Galbraith et al. 2013, Johnson et al. 2016). The climatic and ecological benefits of intact MTFs are potentially threatened by increasing tree mortality due to environmental and biotic changes (Phillips et al. 2009; Lewis et al. 2011, Davidson et al. 2012; Chambers et al. 2013, Erb et al. 2016). Valuable tools for predicting the future of MTF tree mortality are ecosystem and earth-system models (see Glossary; Sperry and Love 2015; Seiler et al. 2015; Levine et al. 2016; Xu et al. 2016). These “next-generation” models have enabled progress on mortality prediction, yet these advances have also revealed multiple questions, particularly regarding MTF tree mortality drivers and mechanisms, that must be addressed to enable accurate prediction (Powell et al. 2013, Thurner et al. 2017). Improving our understanding and model prediction is challenged in part by the enormous variability in mortality temporally, regionally, and within sites according to tree size and other traits (Figure 1).

Accurate prediction of the global climate warming trajectory is challenged by non-mechanistic understanding and simulation of future MTF carbon balance as influenced by tree death (Friedlingstein et al. 2006; Friend et al. 2014). To address this challenge, we describe the state of knowledge of (non-harvest) MTF tree mortality drivers and their associated physiological mechanisms, and investigate the likelihood that these drivers will strengthen in the future. We use empirical and simulation evidence. Through this review, we generate a conceptual framework that provides testable hypotheses regarding the causes, mechanisms, and interactions associated with increasing mortality rates. We briefly investigate factors that may promote survival, and propose a path forward for both empirical and modeling work to better understand the future of MTF tree mortality. Our focus is on intact (primary or old-growth) forests, including aseasonal (wet) and seasonally dry forests, because of their large role in the global carbon cycle (Pan et al. 2011). We are focused on intact forests so that we may investigate if global drivers are associated with mortality, in the absence of direct human intervention. We draw an outer boundary to our geographic scope at the dry margin where forest fires historically occurred. Our scope includes all evidence available from the MTFs in South

America, Africa, and Southeast Asia. We are focused only on mortality; we do not discuss resilience and recovery rates from mortality events, though these are critical questions relative to the terrestrial carbon sink. We use evidence from the extra-tropics when a process appears to be global in nature (e.g. warming impacts on carbon balance) and when tropical evidence is scarce. This ultimately allowed hypothesis generation as to the trends in MTF tree mortality drivers and their mechanisms.

III) Increasing mortality rates in the Amazon Basin

Mortality of individual trees within intact, old-growth forests has been rising during recent decades in the Amazon basin (Figure 2; see Glossary for definitions of mortality rates; unless otherwise specified mortality rate in this manuscript is always defined as % individuals died per total number of live and dead individuals per year), having a significant impact on biomass carbon loss (Figure SI1) and net ecosystem carbon storage (Phillips and Gentry 1994, Phillips et al. 2004, Brien et al. 2015). The trends for the Amazon basin are similar whether plotted as percent mortality rates or biomass mortality (Figure 2 and Figure SI1). These results from 100s of plots across the Amazon are consistent with observed pulse-mortality events in SE Asia (Phillips et al. 2010), and declines in remotely sensed indices (assumed to be correlated with canopy or whole-tree loss) of canopy biomass post-drought in the Amazon (Saatchi et al. 2013) and canopy health in the Congo attributed to drought and warming (Zhou et al. 2014). However, not all tropical forests have exhibited increasing mortality recently (in Panama; Condit et al. 2006, Meakem et al. 2017). The drivers and mechanism(s) underlying this increasing rate of tree death in some areas (while not in others) are currently unknown (Phillips and Gentry 1994; Stephenson et al. 2011; Feldpausch et al. 2016).

At the coarsest level, increasing mortality rates in the Amazon are consistent with observed forest inventory results from old-growth boreal and temperate forests of North America (Figure 2; Luo and Chen 2015). Direct statistical comparison of the lines for the Amazon and for North America is precluded by many limitations (see SI for details), but the similar general trends for the two regions allows for the possibility of similar drivers and mechanisms across North and South America. The Amazon basin has higher mortality rates than North America (Figure 2), which may be expected based on the observed correlation between productivity and turnover at regional (Amazon, Figure 3, and see alternative versions of Figure 3 (Figure SI2A,

B)) and global scales (Phillips and Gentry 1994, Stephenson and van Mantgem 2005; Phillips et al. 2004). We note that an important question arises from Figure (2): is the relationship of mortality rate over time non-linear or linear (our analysis of Brien et al.'s data shows no significant difference between linear and non-linear fits ($p=0.36$; see SI for statistics details). A non-linear pattern is logical because mortality never reaches zero historically, however, a continued non-linear or exponential relationship is also unsustainable. Further discussion of the implications of different statistical fits for Figure (2) are discussed in the SI.

IV) Global and regional mortality drivers and mechanisms

We review mortality drivers that are significant factors in MTFs with the objective of assessing the likelihood that they could already be increasing mortality rates (Figure 2), and ultimately to generate testable hypotheses regarding future mortality rates, their drivers and associated mechanistic processes (Figure 4). We draw upon empirical and simulation evidence of both historical and likely future trends in mortality drivers to aid in generating hypotheses as to the drivers of increasing mortality. In many cases, these expected trajectories are based on limited data (e.g. from the Neotropics) or inferred from uncertain climate forecasts (e.g. wind disturbance) and we have attempted to represent this uncertainty for each trajectory in Figure (4). We review the evidence supporting and conflicting with Figure (4) in the following sections, and include a critical assessment of the data and model limitations. We cannot rank the importance of mortality drivers because there is too little evidence (even at single sites). We focus on tropical evidence throughout our review; however, some drivers (temperature, VPD, and CO_2 in particular) are all rising globally and thus we also use knowledge from the extra-tropics to fill in knowledge gaps when appropriate. While potentially important, nutrient impacts were so poorly covered in the literature that we relegated that text to the SI.

IV.I) Global Driver--Temperature and vapor pressure deficit: Temperature is expected to rise in tropical forests (Figure 4A; Figure 5A-C). MTFs reside in the warmest latitudes on Earth, thus rising temperature will push them into a new temperature regime that has no current analog (Diffenbaugh and Charland 2016). Rising temperature and vapor pressure deficit (VPD) are forcing drivers associated with the multi-decadal increases in tree mortality rates throughout the Americas (Figure 2). There are multiple mechanisms by which rising temperature could cause

rising mortality. First, rising temperature can drive increased respiratory carbon costs via the dependence of respiration on temperature (Clark et al. 2010) and via high-temperature impacts on photosynthetic metabolism, both exacerbating carbon starvation (see Glossary; Figure 4B; Galbraith et al. 2010). Second, rising temperature also causes elevated VPD (Trenberth et al. 2014), forcing greater risk of carbon starvation and hydraulic failure (see Glossary; Figure 4B) via greater stomatal closure and evaporative demand, respectively (McDowell and Allen 2015). Model analyses suggest the impacts of rising VPD on photosynthesis are substantially greater than the impacts of rising temperature *per se* in tropical forests (Lloyd and Farquhar 2008). Rising temperature and VPD can cause a negative carbon balance even at relatively high soil water availability (Zhao et al. 2013). Rising temperatures and VPD may promote biotic attacks (Raffa et al. 2008), though this has not been tested in MTFs. Rising temperature and VPD is also particularly relevant in the mountainous tropics, where mountain tops may limit migration (Feeley et al. 2011, Duque et al. 2015), but also because the range of microhabitats are greater, which could provide refugia under climate change. Impacts of rising temperature and VPD on other mechanisms of mortality are described below (see Figure 4B-E).

IV.II) Global-Regional Driver-Drought: Drought, i.e. precipitation decline that impacts soil moisture, is arguably the best-studied driver of MTF tree mortality. Two critical aspects of drought as a mortality driver are that it episodically occurs everywhere globally, and that the severity of drought extremes is expected to worsen under future conditions (Trenberth et al. 2014; also see Mitigating Factors section below for more details on precipitation forecasts). In particularly wet or anoxic soils the drying may benefit growth and survival, but in many areas this will result in regional increases in mortality (Phillips et al. 2010, Brien et al. 2015, Doughty et al. 2015, Johnson et al. 2016, Powell et al. 2013, Thurner et al. 2017). Droughts happen in MTFs particularly during El Niño events (Ropelewski and Halpert 1987, Ronchail et al. 2002) and periods of warm North Atlantic sea-surface temperatures (Marengo et al. 2011). The most consistent predictions of climate in tropical forests suggest increasing total precipitation (Kitoh et al. 2013, Gloor et al. 2013), but stronger and longer dry seasons over the next century (Boisier et al. 2015; Duffy et al. 2015, Rauscher et al. 2015, Pascale et al. 2016). Due to atmospheric warming (and possibly due to lower relative humidity, see Figure SI3), these future droughts will include higher so-called baseline temperature and VPD than historically

experienced by MTFs (Trenberth et al. 2014, McDowell and Allen 2015), which is the primary driver of the modeled soil drying pan-tropically after 2081 (Figure 5D-G). Thus tropical droughts will be superimposed upon chronically drier soils. In the Amazon basin, dry season length is increasing (Fu et al. 2013), and anomalous droughts occurred in 1997, 2005, 2010 (Marengo et al. 2011), and 2015. In both drought experiments and in observational datasets, the largest trees have disproportionately higher mortality rates under drought stress, with associated large impacts on carbon storage (Nepstad et al. 2007; da Costa et al. 2010, Meir et al. 2015; Bennett et al. 2015; Rowland et al. 2015a; Meakem et al., 2017; Figure 1B). Drought has both positive and negative impacts on the other mortality mechanisms (Figure 4, see text below).

Drought, temperature, and VPD are expected to kill trees alone or via a combination of physiological stress and biotic attack (McDowell et al. 2011). These inter-related mechanisms occur in part via carbon starvation and hydraulic failure (see Glossary; Figure 4B). In particular, sustained periods of severe loss of hydraulic conductivity are a strong predictor of drought mortality in temperate forests (McDowell et al. 2013; Anderegg et al. 2015a; Sperry and Love 2015; Adams et al. 2017), with consistent evidence from the tropics (Rowland 2015a). Carbohydrate status was a strong predictor of mortality in a study of tropical seedlings, with higher carbohydrate content leading to more favorable water status and longer survival (O'Brien et al. 2014).

Moist tropical forests often display paradoxical autotrophic carbon cycle responses to drought. Seasonal and interannual droughts cause greater respiratory carbon loss (Metcalf et al. 2010), lower leaf-level photosynthesis (Doughty et al. 2014), increases in mortality (Phillips et al. 2009, Brien et al. 2015), and reduced regional carbon uptake (Gatti et al. 2014). Nonetheless, droughts sometimes result in stable growth (Doughty et al. 2015, but see Feldpausch et al. 2016 for evidence of decreasing growth) in part via increasing canopy photosynthetic capacity (Clark and Clark 1994, Graham et al. 2003, Saleska et al. 2007, 2016; Huete et al 2006; Brando et al. 2010), flushing of young leaves (Wu et al. 2016), and greater solar radiation (Guan et al. 2015). This paradoxical strategy of prioritizing growth during periods of drought, presumably to compete for light, may accelerate risk of hydraulic failure, carbon starvation, or vulnerability to biotic attack (Doughty et al. 2015). Rowland et al. (2015a) found that both growth and carbohydrate concentrations of trees that survived drought were unchanged relative to control trees, suggesting that survival may either depend on maintenance

of a positive carbon balance, or vice versa, mortality of surrounding trees promotes higher carbon balance in those that survive. Because carbon starvation and hydraulic failure can be induced or exacerbated by myriad drivers, including increases in these processes after fire (Bar et al. 2017), biotic attack (McDowell et al. 2011), and defoliation and shading (Kobe et al. 1997), we hypothesize that carbon starvation and/or hydraulic failure may underlie the mortality resulting from many of the drivers (Figure 4A-E; see hypotheses descriptions below).

IV.III) Global Driver--Carbon Dioxide: Like rising temperature, VPD, and possibly drought, atmospheric CO₂ is rising globally and thus is a candidate driver of the observed increasing mortality rates throughout the America's (Figure 2). But how could rising CO₂ cause elevated mortality rates, when it promotes increased water-use efficiency (Lloyd and Farquhar 2008) and growth? At least two candidate explanations exist. First, at the stand level, rising CO₂ may drive elevated mortality through enhanced growth, which accelerates successional dynamics by driving faster thinning via increased competition for resources (light, water, nutrients). In such a case, the suppressed trees that die experience carbon starvation, hydraulic failure, or biotic attack due to reduced light, water, and nutrients due to increased competition (i.e. the interdependent processes across panels in Figure 4A, B, D, E). Second, rising CO₂ may allow greater growth per individual, thus accelerating the speed at which trees reach large heights, and therefore the rate at which they experience the increased risks of lightning, windthrow, dry-upper canopy environments, and the physiological impacts associated with large size (Nepstad et al. 2007; Bennett et al. 2015; Rowland et al. 2015a). The hypothesis that rising CO₂ may partially drive increasing mortality rates is consistent with 1) the observed mortality rate increase (Figure 2), 2) the relationship between mortality rate and productivity (Figure 3), 3) the relationship between mortality and stand density (Lugo and Scatena 1996), 4) the lag between increases in productivity (first) and then mortality (second) in Amazonia (Brienen et al. 2015), 5) with observed increases in recruitment in Amazonia (Phillips et al. 2004), and 6) the consistent observation that drought-CO₂ studies find little benefit of CO₂ upon survival (reviewed in Allen et al. 2015, but see Liu et al. 2017 for a contrasting model-based result). For these mechanisms to be driving increased mortality, they also must be driving faster stand-level growth but this has only been shown unambiguously for the Amazon basin thus far (Brienen et al. 2015); we lack such tests for African and Asian forests. This idea is not new (Phillips et al. 2004, Stephenson

and van Mantgem 2005, Stephenson et al. 2011), but could be an important driver of increased mortality and thus merits further study.

If either CO₂ (via the enhanced-productivity mechanism), temperature, or VPD drive mortality, then we can expect mortality rates to continue increasing as these drivers are expected to continue rising (IPCC 2014). The remaining mortality drivers discussed below are less certain at the global scale, but evidence exists for them at regional scales.

IV.IV) Regional Driver--Lianas: Lianas (woody vines) are much more common in tropical forests than in temperate or boreal forests (Schnitzer 2005). Lianas reduce productivity and increase mortality of host trees (Figure 1F; Ingwell et al. 2010; van der Heijden et al. 2015, Wright et al. 2015). The total contribution of lianas to tropical tree mortality is difficult to estimate because of wide variation in liana abundance among tropical forests, the relatively small number of studies that have quantified liana influences on tree mortality, differences among studies that make direct comparisons difficult, and the inherent difficulties of quantifying the full impact of lianas on tree mortality. However, Wright et al. (2015) found that 64% of studies had shown liana abundance to be increasing (also see Phillips et al. 2002, Schnitzer and Bongers 2011). Lianas outcompete host trees for resources such as light, water, and nutrients (Johnson et al. 2013), thus they potentially promote both carbon starvation and hydraulic failure of host trees. Furthermore, lianas break limbs and expose fresh wounds for infection by biotic agents. Thus interdependent mechanisms between liana invasion, carbon starvation (e.g. shading), hydraulic failure (e.g. reduced water availability), and biotic agent attack are likely (interactions in Figure 4B-E). Lianas may also increase mortality rates of neighboring uninfested trees, insofar as they increase the rates of treefalls – which can be lethal to smaller neighbors – while competing belowground for water and nutrients (Johnson et al. 2013). Liana abundance tends to increase with dry season length, land use change, and with increasing CO₂ (DeWalt et al. 2015, Granados and Körner 2002, Schnitzer 2014) and thus is expected to increase in the future (Figure 4C).

IV.V) Regional Driver-Fire: Although fires in MTFs are influenced by anthropogenic ignitions, there is a significant role played by climate through drying and increasing fuels (Cochrane 2003, Nepstad et al. 2004, Slik et al. 2010, Brando et al 2014). Droughts increase MTF flammability by reducing understory air and fuel moisture (Ray et al. 2010) and increasing fuel accumulation

from litterfall and mortality (Ray et al. 2005). As a result, forest fires occurring in tropical forests during drought years tend to be larger (Silvestrini et al. 2011, Alencar et al. 2015), more intense, and kill more trees than the ones occurring in non-drought years (Brando et al. 2014). Several lines of evidence suggest that fire seasons in tropical forests have increased over the past few decades (Jolly et al. 2015) resulting in larger (Cochrane and Barber, 2009) and more frequent fires (Alencar et al., 2015). MTF species have few adaptations to resist fires (Barlow et al. 2003, Brando et al. 2012), resulting in even low-intensity understory fires killing a high proportion of the forest community (Barlow et al. 2003, Cochrane and Barber 2009, Slik et al. 2010). Estimates of fire-induced tree mortality rates range from 5% yr⁻¹ to 90% yr⁻¹ (Barlow et al. 2003, Balch et al. 2015, Brando et al. 2016). It is likely that rising temperatures and climate extremes and decreasing surface water content (Figure 5) are increasing forest flammability (Chen et al. 2011). Clear linkages between hydraulic failure and post-fire mortality are now established (Bar et al. 2017) suggesting again that interactions across mechanisms (in this case hydraulic failure and fire) are likely (Figure 4B, C).

IV.VI) Regional Driver--Wind: Wind Convective storms, hurricanes, and typhoons that generate high winds, water logging, and lightning cause tree mortality from individual wind-thrown trees to large blowdown patches (Lugo and Scatena, 1996; Chao et al. 2009, Chambers et al. 2013, Marra et al. 2014). Treefall clusters ranging from individual treefalls to <10 trees per gap represented more than 90% of wind-driven mortality for a Central Amazon landscape (Chambers et al. 2013, consistent with Espirito-Santo 2014a, b). Hurricanes and typhoons also damage forests in coastal and island forests, though these forests are adapted to these events and tend to shed leaves and even branches without complete mortality during wind events (Zimmerman et al. 1994, Yap et al. 2016). Storms are associated in some cases with waterlogging, which promote trees tipping over. Storm-associated lightning also kills trees and damages tree crowns (Magnusson et al. 1996; Yanoviak et al. 2015), but has been little studied in MTFs even though lightning frequencies are higher in the tropics (Christian et al. 2003). No study has yet determined if wind-associated mortality has a latitudinal trend at the global scale, though there is a latitudinal trend in average wind speed, average wind speed declines towards the tropics (<http://globalwindatlas.com/datasets.html>) and equatorial regions (≤ 10 degrees from the equator) rarely experience hurricanes/typhoons. Extreme storm events are expected to

become stronger and more frequent with climate warming (Emanuel 2013; IPCC 2014, see SI Figure SI4) with warming-driven increases in atmospheric latent heat, indicating a shift toward more intense wind disturbance regimes in MTFs (Figure 4D).

IV.VII) *Regional Driver-Biotic agents:* Pathogens, insects, and other biotic agents contribute to tree mortality (Coley and Barone 1996) and play a strong role in structuring tropical forests (Mangan et al. 2010, Coley and Kursar 2014). While only rarely studied, heart rot is associated with >50% of stems in a forest in Borneo, and may be strongly associated with susceptibility to wind events that cause loss of branches, stem breakage, or windthrow (Heineman et al. 2015). Far less is known about tropical outbreaks of biotic agents than temperate outbreaks leading to unclear expectations of their response to future climate (Figure 4D), due in part to the great diversity of species that kill trees (Dyer et al. 2012) and the historic focus on defoliators that often do not kill trees (Anderegg et al. 2015b). However, attack by insects was greater in a drought experiment in the Amazon (Brando et al. 2006) and tends to follow droughts (Anderegg et al. 2015b). Biotic agents often cause widespread tree mortality events in the temperate and boreal zones (Kautz et al. 2017), but die-offs of the magnitude observed in low-diversity forests (Breshears et al. 2005) have not been observed in tropical forests. The largest mortality rates observed in moist tropical forests rarely exceed 5% (Figure SI1), whereas mortality events exceeding 90% of individuals lost have occurred in the extratropics (Breshears et al. 2005), generally the result of a drought-facilitated insect (e.g. bark beetle) outbreak upon single or multiple species. The relatively low rates of mortality in MTFs (compared to the extra-tropics) may be due to the high species diversity and the relatively high specificity of biotic agent-host tree relationships, coupled to the asynchronous timing of outbreaks of biotic agents (Dyer et al. 2007; Coley and Kursar 2014). Alternatively, the rate of biotic-attack driven mortality may be higher but less detectable in the tropics than in the extra-tropics. Thus while biotic agents are clearly important mortality drivers in MTFs, their historical or expected future trends in attack rates are poorly constrained (Figure 4D).

IV.VIII) *Regional Driver--Shading:* Shading in light-limited MTFs is an expected driver of mortality (Wright et al. 2010, Ruger et al. 2011) and has been associated with carbon starvation in four species of angiosperms (Kobe 1997). The dichotomy between the low light environment

and the high light environment when gaps form has had a distinct impact on evolutionary strategy of species (Richards 1952). Slow-growing, shade-tolerant trees tend to live longer than fast-growing, shade-intolerant trees (Condit et al. 1995, Wright et al. 2010; Figure 1D). Shading is presumed to be the dominant driver of the high mortality rates of seedlings and understory plants (Figure 1A, Panama example), however the mechanisms of the interactions between shade, herbivory, biotic agents, and the physiological mechanisms of carbon starvation and hydraulic failure (O'Brien et al. 2014) within the ultimate mortality process is poorly known. Solar radiation is expected to increase in much of the tropics (Collins et al. 2013), and rising temperature and VPD would act to further reduce shading by inducing mortality (or lower leaf area) of competing vegetation. In contrast, the competitive dynamics that drive mortality via shading may be speeding up due to CO₂ induced increased productivity (Brienen et al. 2015) and higher leaf area. Thus, there is large uncertainty in the trajectory of shading in the future (Figure 4E).

IV.IX) Summary – mortality drivers

In summary, amongst the identified mortality drivers in tropical forests, most appear to be increasing in potential or frequency, thus there is reasonable evidence to conclude that risks to continued increases in tree mortality within moist-tropical forests are likely. Temperature, VPD, fire, wind, biotic agents, lianas, and potentially CO₂-induced thinning and accelerated height growth (Figure 3) may all possibly increase under future climate change (Figure, 4). However, the lack of knowledge of the relative impacts and interactions of each process on MTF tree mortality, and inadequate evidence of their trajectories (particularly for competition) make determination of the relative causes of rising mortality rates (Figure 2) a challenge both historically and in the future.

V) On the coupling of mortality drivers and mechanisms

Mortality drivers and mortality mechanisms (see Glossary for definitions) are coupled through a chain of events, starting from an initial forcing variable that promotes an increase in a mortality driver (e.g. rising CO₂ forces rising temperature), and the mortality driver subsequently impacting plants via structural (e.g. windthrow) or physiological mechanisms (e.g. liana-shading reducing photosynthesis; Figure 4). Understanding these linkages is valuable both from a

fundamental knowledge perspective and for advancing mechanistic-mortality simulation within newer ESMs. We previously explained the linkage between carbon starvation, hydraulic failure, and temperature, VPD and drought, and now hypothesize on how these mechanisms are tied to the other mortality drivers (Figure 4).

Fires and wind-events can destroy entire trees via simple structural breakage. For the other mortality drivers, we propose that drivers kill trees via the mechanisms of carbon starvation (and phloem failure) and hydraulic failure (see Glossary for definitions). Carbon starvation should be promoted by increased shade from neighboring trees or lianas, and can be further exacerbated if liana's girdle the phloem. Defoliation from wind and insects promotes carbon starvation if sufficient canopy is removed, though such disturbances may need to be repeated in high frequency to sufficiently deplete stored carbohydrates (Wurth and Korner 2005). Biotic agents may successfully invade trees that have low carbohydrates from the carbon starvation process and low sap pressure (Lorio and Hodges 1968). Hydraulic failure may be promoted by increased competition for soil water such as from lianas, and fire promotes hydraulic failure in partially burned trees (Michaletz et al. 2012) thus resulting in greater death than the consumed stems alone. The carbon starvation and hydraulic failure framework has had a growing impact on ESMs (Fisher et al. 2010; 2015; McDowell et al. 2013) because it is logical and consistent with available data, however, extending it (including validation) to include the interactions with lianas, wind, fire, shade, and other drivers has yet to be attempted. Whether representing carbon starvation and hydraulic failure associated with the myriad mortality drivers will improve model predictions over simpler empirical functions is an emergent question as we begin to uncover mechanisms.

V) Mitigating factors that may promote future survival

There are potential mitigating factors that may promote survival of trees in MTFs that should be considered. The three most obvious mitigating factors are species diversity (Poorter et al. 2015), rising CO₂ impacts on carbon and water relations (Keenan et al. 2016), and the potential of increasing mean annual precipitation (Figure SI5).

Higher species richness and hence physiological traits are expected to reduce vulnerability to large-scale mortality events (Mori et al. 2013). Empirical data from tropical forests suggests higher diversity does beget greater resistance to drought in terms of individual

mortality rates (Williamson et al. 2000; Fauset et al. 2012) and sometimes carbon storage (Poorter et al. 2015, but see Sullivan et al. 2017). The mechanisms by which diversity promotes resistance (ability to withstand change) and resilience (ability to recover) are thought to lie in the greater capacity of the forest community to tolerate new conditions due to a wider range of traits that enable survival (e.g. hydraulic traits that promote drought tolerance; Christoffersen et al. 2016; Powell et al. 2017). Evidence on the role of diversity in global patterns of mortality comes from comparison of rates of drought-induced death in the moist tropics, where mortality rates (on an individual basis) are rarely above 5% in inventory plots even after droughts (Figure SI1B) and only up to 15% in drought experiments (Nepstad et al. 2007, Rowland et al. 2015a), versus the temperate zone where mortality rates can exceed >90% (Breshears et al. 2005; Plaut et al. 2012).

As reviewed earlier, elevated CO₂ benefits water-use efficiency (Ehrlinger and Cerling 1995, Lloyd and Farquahar 2008), but the degree to which this results in changed growth at the individual tree level remains disputed (van der Sleen et al. 2015, Brien et al. 2016). Enhanced growth should result in less risk of mortality of the trees that are rapidly growing (Chao et al. 2008), as should enhanced water-use efficiency through reducing the risk of both hydraulic failure and carbon starvation. However, CO₂ manipulation studies that imposed drought and killed trees rarely found any effect of CO₂ on survival (all greenhouse studies; reviewed in Allen et al. 2015). It remains a large question what the impact of CO₂ is on moist-tropical tree mortality and this introduces uncertainty into the associated drivers (Figure 4).

Increasing mean annual precipitation may occur in some tropical regions (Figure SI5). This would act to only partially buffer the large increase in evaporative demand due to temperature (Figure 5A), which results in significant reductions in soil moisture (Figure 5B) based on the Coupled-Model Intercomparison Study (CMIP5, Collins et al. 2013). As reviewed earlier, the occurrence of droughts that are warmer than previously will increase, thus their impact will be more severe (Trenberth et al. 2014). There is some prediction of shifts to longer drought lengths (Boisier et al. 2015; Duffy et al. 2015, Rauscher et al. 2015, Pascale et al. 2016). Note that increasing precipitation, when it does occur, also results in greater shade, more soil anoxia, and greater windthrow, so it is not clear that there will be much net benefit of increasing precipitation on survival of moist-tropical forest trees.

VI) The state of ESM simulations of moist tropical tree mortality

ESM's are the required tool to predict moist-tropical tree mortality pan-tropically. However, many ESM processes, including those relevant to mortality, draw upon ecosystem- and individual-plant scale models in part because they provide mechanistic simulation capabilities at appropriate scales (e.g. the individual plant). As discussed above, while there is evidence of increasing likelihood of mortality drivers, we still need substantially more data on these processes in order to understand them sufficiently to model them. As a result, many of the mortality drivers and mechanisms discussed here (Figures 1-4) are not represented in ESMs, and thus accurate simulation of the future mortality-related carbon flux requires process development. Before discussing the next steps in empirical and model developments, we briefly review the state of ESM simulations of mortality in moist-tropical forests.

Most tropical ESM projections highlight the interaction between the fertilization impacts of rising CO₂ and the deleterious impacts of increasing drought and heat stress (Cox et al. 2004; Huntingford et al. 2008; Fisher et al. 2010; Rowland et al. 2015b). However, many earlier-generation ESMs simply assume a fixed mortality rate (often called background mortality, see Glossary), leading to a growth-only driven estimate of forest carbon fluxes and stocks (i.e. they cannot capture the trends in Figure 2; de Almeida Castanho et al., 2016; Johnson et al. 2016; see Table 1 within McDowell et al. 2011 for a brief summary of mortality mechanisms in ESMs). This is a significant problem because ESMs must simulate mortality sufficiently well to properly predict ecosystem biomass (Galbraith et al. 2013; Johnson et al. 2016), particularly if mortality drivers are changing (Figure 4).

Among the newer generation of ESMs, two representations of mortality are common. The first is shifting from one plant functional type (PFT) to another (representative of mortality and regeneration by a new type) based on climate envelopes (Sitch et al., 2003). The second is the use of constant biomass residence times (see Kucharik et al., 2006), which is tantamount to assuming "senescence" mortality, in which a genetically predisposed age threshold is used. Both of these approaches risk over-simplification. Climate envelopes do not capture spatial variability such as with different climates, species, or topography, and may not be realistic in a future, warmer, higher CO₂ world. Age-driven mortality, while it may capture the statistical odds of dying from pathogen infestation, wind, or lightning, is not mechanistically representative (Mencuccini et al. 2005) and may thus also fail under a novel climate.

A more sophisticated yet common approach to simulate tree mortality in ESMs is the use of growth efficiency, in which a PFT is replaced if its stemwood growth per individual leaf area is below a threshold (McDowell et al. 2011). The low growth-efficiency approach is mechanistic and supported because trees that die tend to grow more slowly (per unit leaf area) than those that live (Chao et al. 2008, McDowell et al. 2008, Cailleret et al. 2016) and because growth is intimately tied to carbon starvation (McDowell 2011). Furthermore, the growth-efficiency approach responds to most if not all climate drivers that limit growth, including CO₂, light limitation, drought, and VPD. Next-generation approaches that are under current or planned development, as well as new ideas on ESM developments that have not yet been attempted, are discussed in the ensuing sections on specific ESM development needs.

VII) Next steps

There are numerous hypotheses regarding the possibility of increasing future MTF mortality rates (e.g. continuation of trends in Figure 2) that revolve around the dependence of mortality process changes, and subsequent mortality rate changes, on chronic or punctuated changes in mortality drivers (Figure 4). We outline our highest level hypotheses here:

- 1) MTF mortality rates are increasing linearly and will continue under projected climate change (Figure 2);
- 2) mechanisms of mortality e.g. lianas, fire, biotic agents, wind, competition, and shade, are increasing
- 3) with the exception of death from direct physical destruction (e.g. windthrow or intense fire), mortality involve a cascade of impacts from driver (Figure 4A) through a mechanism (Figure 4C-E) to a physiological death process (Figure 4B);
- 4) uncertainty can be reduced through quantifying the primary mechanisms and processes underlying rising mortality rates in MTFs.

Many sub-hypotheses have been previously outlined and will be expanded upon below, but all revolve around the trajectories and interactions between expected drivers, their mechanisms, and physiological end points (Figure 4).

VII.I) Observations: We do not know the relative importance of the various drivers of MTF mortality (Figures 1-4) nor do we have sufficient confidence in the trajectory of these mortality

drivers in the future to make rigorous predictions (Figure 4). Quantifying the various mortality mechanisms in MTFs is limited by a scarcity of temporal and spatial data sufficient to overcome the high signal-to-noise ratio inherent in field observations of plant mortality. Long-term and high-temporal frequency observations (e.g. annual) at the plot-level are essential to reveal the long-term spatial and temporal patterns of mortality in relation to climate dynamics. Plot networks, although challenging to run, are arguably the lowest cost, highest impact investment one could make to refine the uncertainty in moist-tropical mortality drivers. Plot networks provide information regarding the dynamics of growth and death in response to droughts (Condit et al. 1995, Phillips et al. 2009; Brien et al. 2015, Anderson-Teixeira et al. 2015), and with appropriate measurements, they can unveil mechanisms driving mortality (Doughty et al. 2015). A relatively low-cost addition to inventory networks could be assessment of the “modes” of death (snapped, died standing, windthrow, presence of rot, etc), determination fraction of crown shaded (by neighbors or lianas), and dendrometer measurements prior to death. Plot-level work can in some cases include tree rings, even for tropical trees (Schöngart et al. 2006, van der Sleen et al. 2015; Brien et al. 2016), which can provide proxy measurements of physiology proceeding death (Gaylord et al. 2015). Similarly, remotely-sensed data provide unparalleled spatial coverage of drought impacts, such as the long-term decline in canopy health associated with declining precipitation and increasing temperature in the Congo Basin (Zhou et al. 2014) and the sustained loss of biomass observed post-drought in the Amazon (Saatchi et al. 2013). A key step is validation of remote sensing estimates of mortality against ground-based data such as mortality rates, leaf area, canopy height, and canopy biomass and correlations of remotely sensed indices of dying and surviving trees at the crown scales e.g. using high-resolution (<10m) satellite products now available (McDowell et al. 2015).

VII.II) Experiments: Cause-and-effect experiments that manipulate mortality drivers (Meir et al. 2015, van der Heijden et al. 2015) are valuable because they can reveal the mechanisms underlying mortality, and can be employed for model evaluation under novel climate conditions. The few moist tropical drought experiments (Nepstad et al. 2007, Moser et al. 2014, Rowland et al. 2015a, Meir et al. 2015), cannot be representative of the diverse MTFs and thus experiments replicated across a broad range of soils, topographic relief, and proximity to groundwater (Nobre et al., 2011) are needed. Replication of such experiments across a wider range of sites in the

moist-tropics could be achieved economically if the measurement intensity was low. However, in addition to replication, some of the next generation experiments must address the multi-factorial climate changes expected in the future e.g. low precipitation and elevated CO₂ or rising temperature (and associated rising VPD), and should push drought to extreme levels to understand acute impacts and threshold responses (Knapp et al. 2016) including mortality. Otherwise, such experiments manipulate only one of the many variables that are changing, and thus determining the net effects under future climate scenarios is challenged. Multi-factorial and replicated experiments have not been conducted in mature tropical forests for financial, technical, and logistical reasons. The most challenging aspects of manipulative experiments are their inability to control all environmental conditions, and their minimal replication relative to the hyper-diversity of tree species in MTFs.

VII.III) *ESM Demographics:* To allow simulation of competition, shading, lianas, and size-dependence of mortality as they may change over time (Figure 4), ESMs should represent demographic heterogeneity in vegetation (horizontal and vertical size variation, Moorcroft et al. 2001; Fisher et al. 2015; Levine et al. 2016). Big-leaf (no demography) model simulations predict trees fail to die (Powell et al. 2013) or die more often and faster than is observed (Galbraith et al. 2010; Poulter et al. 2010), whereas the addition of demographic variation in size and environment results in more realistic, gradual mortality (Powell et al. 2013; Levine et al. 2016). Simulating demography allows more realistic spatial heterogeneity in resource capture and loss and thus better simulations of mortality against observations, for example, prediction of taller trees dying in a drought experiment (Longo 2013).

VII.IV) *ESM drought, temperature, VPD and CO₂:* Given that mortality is downstream of the majority of other physiological processes (assimilation, respiration, allocation), predictions are sensitive to assumptions about photosynthesis, respiration, carbon allocation, and carbon storage (Fisher et al. 2010), all of which are heavily influenced by plant hydraulics (Christoffersen et al. 2016; see text below on hydraulic modeling limitations and developments) and so predictions tend to be extremely divergent among models (Galbraith et al. 2010, Huntingford et al. 2013). To improve accuracy under non-linear changes (and complex interactions) of future drought, temperature, VPD, CO₂, and hopefully someday wind, fire, and lianas, next-generation models are now including more realism such as carbon starvation and hydraulic failure (Fisher et al.

2010, 2015; McDowell et al. 2013; Sperry et al. 2016; Xu et al. 2016), though evaluation in MTFs is needed. Simulating these mortality mechanisms requires representing water transport, xylem embolism, photosynthesis, and carbon storage accurately.

Inclusion of plant hydraulics allows more realistic simulation of mortality (McDowell et al. 2013, Anderegg et al. 2015a) and photosynthesis (Bonan et al. 2014). Thus simulating plant hydraulics allows more accurate representation of both the risk of hydraulic failure and the likelihood of carbon starvation under changing climate, and of the interactions of these processes with external drivers such as lianas, shading, biotic agents, wind, and climate. Most land components of ESMs model plant response to drought as a function of the vertical profile of prescribed fine root biomass ('root fraction' in models) and soil moisture, and collapse these two profiles into a single non-dimensional ('beta') multiplier [0,1] that is applied to Ball-Berry stomatal parameters or to carbon assimilation (Sitch et al. 2003, Krinner et al. 2005, Kucharik et al. 2006, Oleson et al., 2010). Three main reasons exist why this approach is insufficient for modeling tropical forest hydraulic and subsequent carbon assimilation responses to reductions in moisture. First, these models poorly capture observed experimentally-induced patterns of mortality (Powell et al., 2013, Joetzjer et al. 2014) in contrast to site-specific models that include plant hydraulics (Williams et al. 1998, Fisher et al., 2006, 2007). This model-observation mismatch is due in part to the 'beta' approach: because all trees' drought response is considered equivalent and shares the same threshold response, causing an all-or-nothing response to drought. Second, current approaches lack the ability to model a well-documented negative interactive effect of soil moisture and VPD (Sperry and Love 2015, Sperry et al. 2016), which plays an important role in regulating tree response to typical droughts. Finally, a wealth of knowledge regarding plant hydraulic traits that govern how tropical trees transport and use water under a range of moisture conditions has been synthesized in multiple databases that quantify inter- and intra-specific variation (Bartlett et al., 2012, 2014, 2016; Choat et al., 2012, Gleason et al., 2016, Christoffersen et al., 2016, Wolfe et al. 2016). While the typical argument against increasing model process complexity usually states that a host of unknown parameters are introduced, the case of plant hydraulics represents the opposite: parameter central tendencies, ranges, and variances are already known but most current model structures are incapable of exploiting this information. Inclusion of biophysically-based representations of water acquisition, transport, and use holds great promise for increasing the realism of tropical forest

drought and mortality responses (see an example approach for future ESM hydraulic development in the supplemental information).

Carbon starvation is sensitive to shade, temperature, VPD, and CO₂ (Figure 4; reviewed by McDowell et al. 2011) among other factors. In practice, carbon starvation mortality is simulated as a response to nonstructural carbohydrate stores; i.e., trees die when nonstructural carbohydrate stores reach zero (Weng et al. 2015), or when carbon storage is less than leaf biomass carbon (Fisher et al. 2010), though these thresholds are arbitrary and more work is required to determine if a universal threshold exists under field conditions (Adams et al 2017). The accuracy of carbohydrate simulations can be high (e.g. McDowell et al. 2013), but observations of carbohydrate content at death are required to tune models to simulate mortality via carbon starvation, because the carbohydrate concentrations at death are variable (Adams et al. 2017) and because carbohydrate results vary between labs/studies (Quentin et al. 2015). Furthermore, the role of carbon in mortality remains in question, therefore carbon starvation by itself may not be the appropriate mechanism to simulate tree death (Rowland et al. 2015a), but rather an interdependency of carbon starvation and hydraulic failure, and linkages to phloem failure may be required to improve model simulations during drought or under low light (O'Brien et al. 2014, Sevanto et al. 2014, Mencuccini et al. 2015; Adams et al. 2017).

VII.V) *ESM trait-based modeling in the diverse moist tropics:* Modeling the myriad set of mortality drivers and mechanisms (Figure 4) is challenging as it requires identification and incorporation of the trade-off and coordination among different traits targeted for different survival strategies (Fisher et al. 2015). This is a particularly important issue in the particularly diverse tropics, where the variety of species and thus traits are greatest, but are represented by only a limited number of PFTs used to model MTFs i.e. evergreen vs. deciduous trees. Next generation models are moving towards becoming trait-enabled such that trait-trade-offs facilitate simulation of diversity impacts on carbon and water balance of forests (Sakschewski et al. 2016). Data to parameterize these models is becoming available at the global scale, with discovery of quantitative relationships among plant traits (Wright et al. 2004, Christoffersen et al. 2016), the inter- and intra-specific and biogeographical components to their variation (Anderegg 2015c), the number of independent axes of trait variation in forest communities (Wright et al. 2007, Baraloto et al. 2010, Reich 2014), and relationships of plant traits to tree mortality (Wright et al.

2010). For example, many parameters required for simulating plant hydraulics (such as pressure-volume relationships) can be estimated from traits such as wood density (Christoffersen et al. 2016). This understanding informs us how models can represent new and flexible PFT definitions (Pavlick et al., 2013, Verheijen et al. 2013, Harper et al. 2016; Powell et al. 2017), which is a critical prerequisite for developing modeling capability to represent ecological sorting mediated by plant traits (i.e., trait-mediated environmental filtering *sensu* Sommer et al. 2014). It is important for next-generation ESMs to predict shifts in trait distributions through time (Scheiter, Langan & Higgins 2013) because of mounting evidence showing that key aspects of ecosystem-level properties (e.g., C storage, overall resiliency) depend on the functional community composition (Fauset et al., 2012). A critical challenge, however, is for us to better understand what traits, their trade-offs, and their plasticity (Lloyd et al. 2010), result in tolerance or susceptibility to mortality drivers (Figure 4).

VII.VI) *ESM Lianas*: No ESMs have yet attempted to explicitly represent lianas (Verbeeck & Kearsley 2016). The empirical knowledge base for modeling lianas is incomplete, but our existing knowledge regarding the role of gaps, CO₂, and drought on liana abundance can provide some simulation potential for liana succession. With demographic ESMs it may be possible to simulate the succession and impacts of liana's on upper-canopy trees through shading and breakage, particularly in gaps. Trait-enabled hydraulic models will be able to simulate the high rates of soil water acquisition by lianas (Johnson et al. 2013) and subsequent impacts on host tree water availability. For mortality mechanisms, lianas likely impact hydraulic failure through drawing down soil moisture via their high transpiration rates (Chen et al. 2015), and carbon starvation via shading, but determining the fraction of host-crown shaded, and impacts on water consumption, are required to inform model mechanism.

VII.VII) *ESM Fire*: Most ESMs include representations of fire, but the majority of these models are parameterized from limited studies in boreal and temperate regions, and their applicability to tropical systems is largely unknown (Hantson et al 2016). Improvements in the simulation of fires for the tropical forests should focus on 1) mechanism-scale validation of fire spread and tree mortality simulations against fire experiment data, 2) tests of how fire-vegetation interactions are simulated at stand-to-ecosystem scales, and 3) developments that focus on the landscape-scale

determinants of fire durations, maximum fire extent, the geographical spread of ignition events and interactions with human activity. The latter problem in particular poses significant issues concerned with how to attribute patterns observed through remote sensing to variation in different processes (ignition, suppression, fragmentation), and with predictive models of interactions with human behaviors. Increasing abundance of regional and global fire remote sensing products (Alencar et al. 2015, Bloom et al. 2015) allows at least the possibility of better landscape-scale calibration of the higher-level features of such models, while more robust testing of physical models of fire spread should increase confidence in our ability to predict responses to altered climatic drivers in future scenarios.

VII.VIII) *ESM Biotic agents:* Most ESMs have not simulated biotic attacks (insects and pathogens; but see Dietze and Matthes 2014, Landry et al. 2016) but a path forward can be derived from a few key observations. Insect outbreaks often occur after droughts in the moist tropics (Anderegg et al. 2015b), exhibit a correlation between host tree defense and outbreak success in both temperate (Herms and Mattson 1992, Raffa et al. 2008) and tropical regions (Dyer et al. 2007), and outbreaks (i.e. widespread attacks on one or more species) decline with increasing diversity at the global scale (Jactel and Brockerhoff 2007). Less is known about the processes driving biotic agents such as heartrot and rootrot, but we may presume that infection by these agents is similar in physiological regulation to that of insects (see McDowell et al. 2011). Thus an initial ESM approach could be to simulate defense (perhaps using available carbon as a surrogate) and assume (for now) that biotic agents are ubiquitous in presence. However, in addition to predisposition by plant stress, outbreaks of tropical tree-killing insects are also more likely after other types of disturbances that open the canopy and increase the abundance of light, new foliage, and juvenile trees (Dyer et al. 2012), which suggests that the dynamics of canopy gap formation in demographic models may be used for outbreak initiation. Although these bottom-up controls by plant defenses and stand structure play a role in outbreaks of tropical tree killing insects, top-down predator control appears particularly important in the tropics in constraining the magnitude of outbreaks (Van Bael et al. 2004). Thus an idealized model might include a function associated with host tree defense capability, host-tree abundance (Dyer et al. 2012), forest structure (Dyer et al. 2012), insect thermal optima (Goodsman et al. in revision), and top-down insect predator abundance, all influenced by environment.

VII.IX) *ESM Wind*: Arguably the hardest ESM challenge is to downscale maximum wind speeds from atmospheric models that simulate average wind speeds over the scale of individual grid cells (e.g. Figure SI4) and are formulated using a hydrostatic approximation that prevents explicit representation of processes that generate high wind extremes. At the canopy-scale, the ability to model either loss of foliage, loss of major branches, snapped-stems, standing dead stems, or an uprooted tree is valuable for capturing recovery processes, gap light dynamics, and carbon cycling from wind mortality (Holm et al. 2017), which can be most aptly simulated in demographic models. Opportunities to further improve predictions of wind mortality lie in representing abiotic and biotic conditions (e.g., soil conditions, prior exposure to stress, presence of heartrot) that enhance vulnerability to wind, traits that confer susceptibility or resistance to wind, and the wind fields that can topple canopy trees (Ribeiro et al. 2016).

VIII) Conclusions

Many of the drivers of MTF tree mortality appear to be increasing (Figure 4, though with large uncertainties), thus there is some confidence that mortality rates may increase over time. These mortality drivers may include productivity-driven thinning and increase in height growth, rising temperature and VPD, increasing frequency and severity of droughts, increasing liana competition, fire, wind disturbance, and biotic attacks. Determining the relative importance of these drivers is critical to enable mechanistic prediction of future mortality. Simulating future tropical forest mortality under climate-change is daunting due to this lack of knowledge coupled with the complexity of processes in hyper-diverse tropical systems. Some model mechanisms require improvement, such as including refined hydraulics and demographics, whereas other model processes have yet to be included, such as wind, insects, and liana competition. Model structures that include demographic representation and represent the diversity of physiological traits should provide a useful foundation for rapid model development, but such development must progress hand-in-hand with increasing empirical knowledge of the key processes that regulate tropical forest mortality under climate change.

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For Peer Review

Box 1. Glossary

Background mortality: also considered a fixed mortality rate (e.g. % year⁻¹, carbon m⁻² yr⁻¹) in models and referred to as such in this manuscript; this is the theoretically stable mortality rate under a non-changing environment.

Biotic agents: insects, fungi, and other pathogens that attack and sometimes kill trees directly or by weakening them (e.g. defoliation, or rot impacts on wind resistance).

Carbon starvation: the *process* by which limited carbon uptake (e.g. due to stomatal closure, shade, or leaf area loss to wind damage) relative to carbon demand (e.g. growth, respiration, defense) results in a decline in carbon-driven metabolism, hydraulic repair, or ability to defend against pests, and ultimately promotes mortality (McDowell et al. 2011).

Earth System Model (ESM): models designed to simulate the coupled influences and feedbacks of climate, land and ocean. Land surface models operate within ESMs.

Hydraulic failure: mortality via dehydration; often associated with prolonged periods of xylem conductivity loss >60% in field studies (McDowell et al. 2013).

Lianas: woody plants that utilize free-standing hosts to support their weight as they grow into the canopy. Lianas are typically aggressive consumers of light, water, and nutrients.

Moist-tropical forests (MTFs): forests with mean annual precipitation > 1500 mm, including both aseasonal and seasonal precipitation regimes (e.g. with a dry season < 100mm/month for five months or less; Vitousek and Sanford 1986).

Mortality drivers: factors that when they experience a directional change so do mortality rates. Examples include decreasing precipitation, increasing temperature, and increasing biotic attack.

Mortality mechanisms: mortality drivers cause changes in mechanisms that lead to mortality, such as altering plant structure (e.g. via windthrow, fire) or physiology (e.g. shade-induced carbon starvation, drought-induced hydraulic failure).

Mortality rate: Can be defined using many units, typically % yr⁻¹ (number of trees died per number of total individuals live and dead per year) or in units of basal area (m² basal area died/m² of total stems/year) or biomass (kg C died/kg C standing biomass/year). Corrections for biomass weighting, non-balanced plot sizes or sampling periods over time and space are often employed when calculating mortality rates from inventory data. See Supplemental Information for equations.

Figure 1. Axes of variability in tropical tree mortality. A) Mortality rate [as $\log(\text{initial number}) - \log(\text{number survivors})/(\text{years})$] versus stem diameter in Pasoh, Malaysia and Barro Colorado, Panama (bars are 95% CI, no major droughts during censuses); this highlights that both negative and positive mortality rates as a function of diameter can be found (from Muller-Landau et al. 2006). B) Mortality rates (# individuals died per number of total individuals per year; all subsequent figures use this calculation; see Glossary and SI for information on mortality rate calculations) plotted as the ratio of mortality rate during drought relative to a control period across a range of stem diameters for 12 sites across the tropics (symbols represent different sites), showing the clear pattern size-mortality relationships *during droughts* (from Bennett et al. 2015). C) The mortality rates in forests in Borneo and the Amazon measured post-drought, highlighting regional differences (from Phillips et al. 2010). D) Mortality rate versus life-history strategy in Barro Colorado, Panama, highlighting the role of successional strategy on long-term mortality rates (from Condit et al. 1995). E) Mortality rate versus wood density in Barro Colorado, Panama, highlighting a significant but weak relationship ($p < 0.05$; from Wright et al. 2010). F) Mortality rate as a function of liana cover class in Pasoh, Malaysia, highlighting the influence of lianas on mortality. Liana cover class 0 indicates no lianas, 1 indicates up to 25% of the crown covered by lianas, 2 = 26-50%, 3 = 51-75%, and 4 = 76-100% (from Wright et al. 2015).

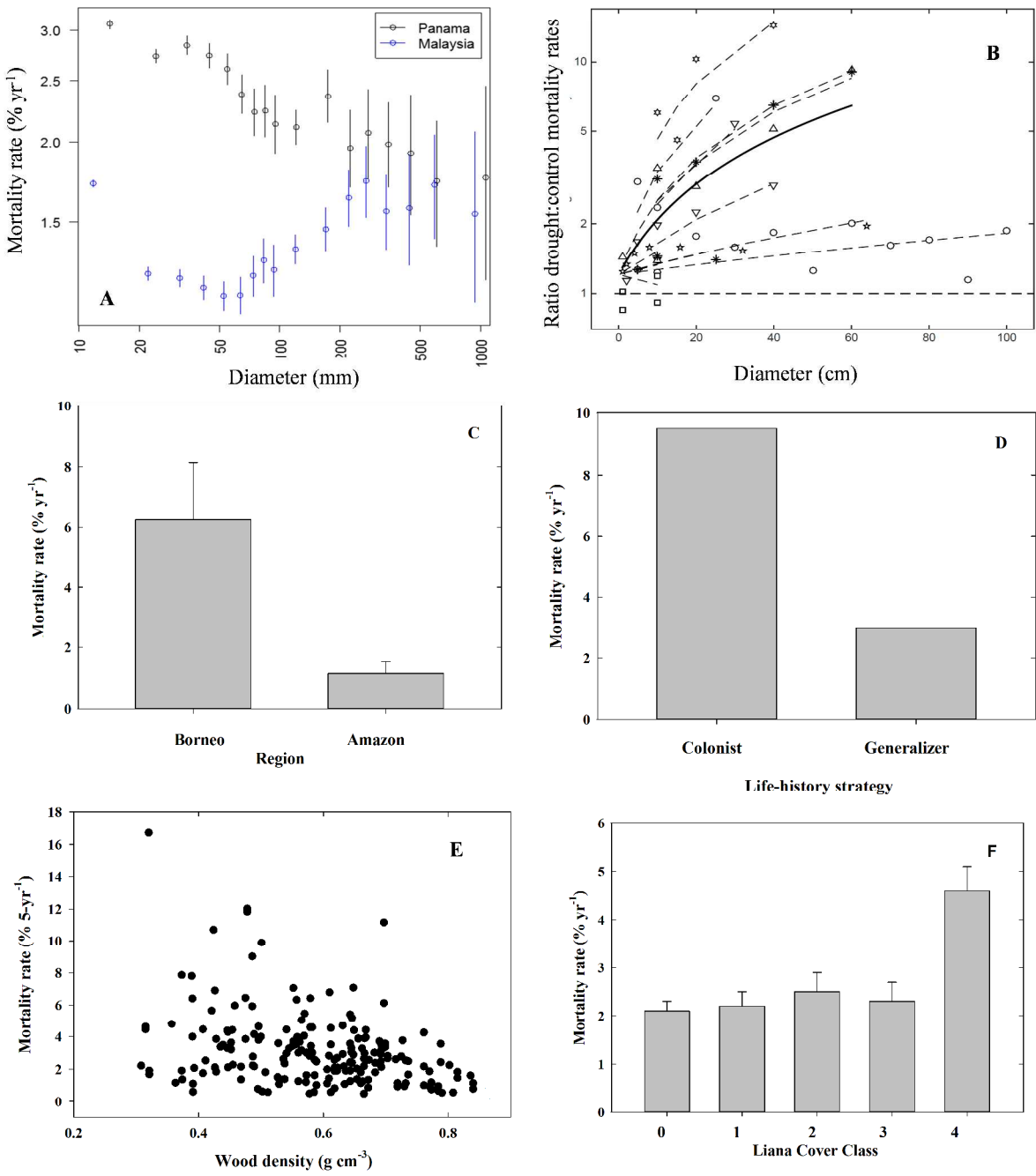
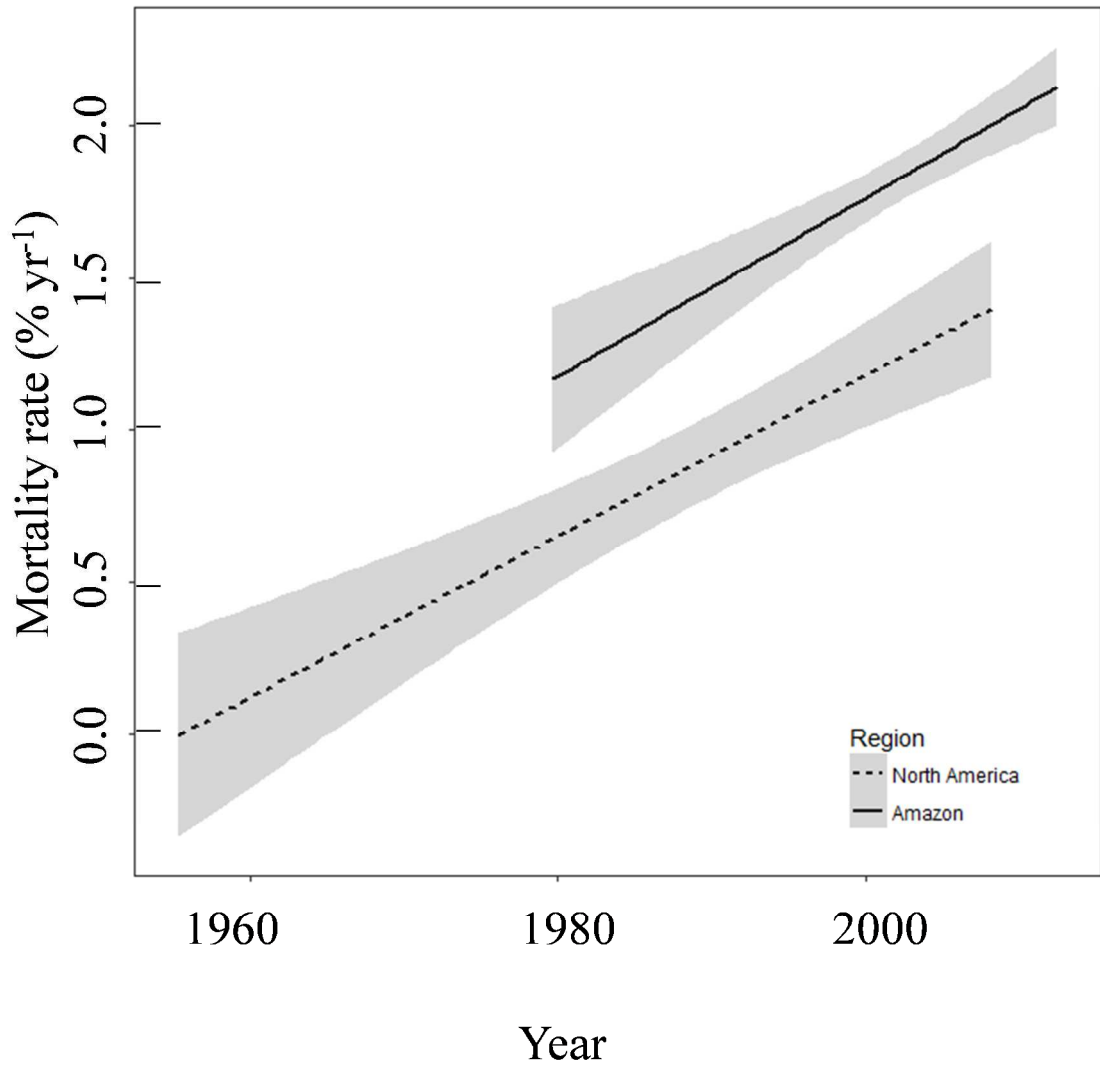
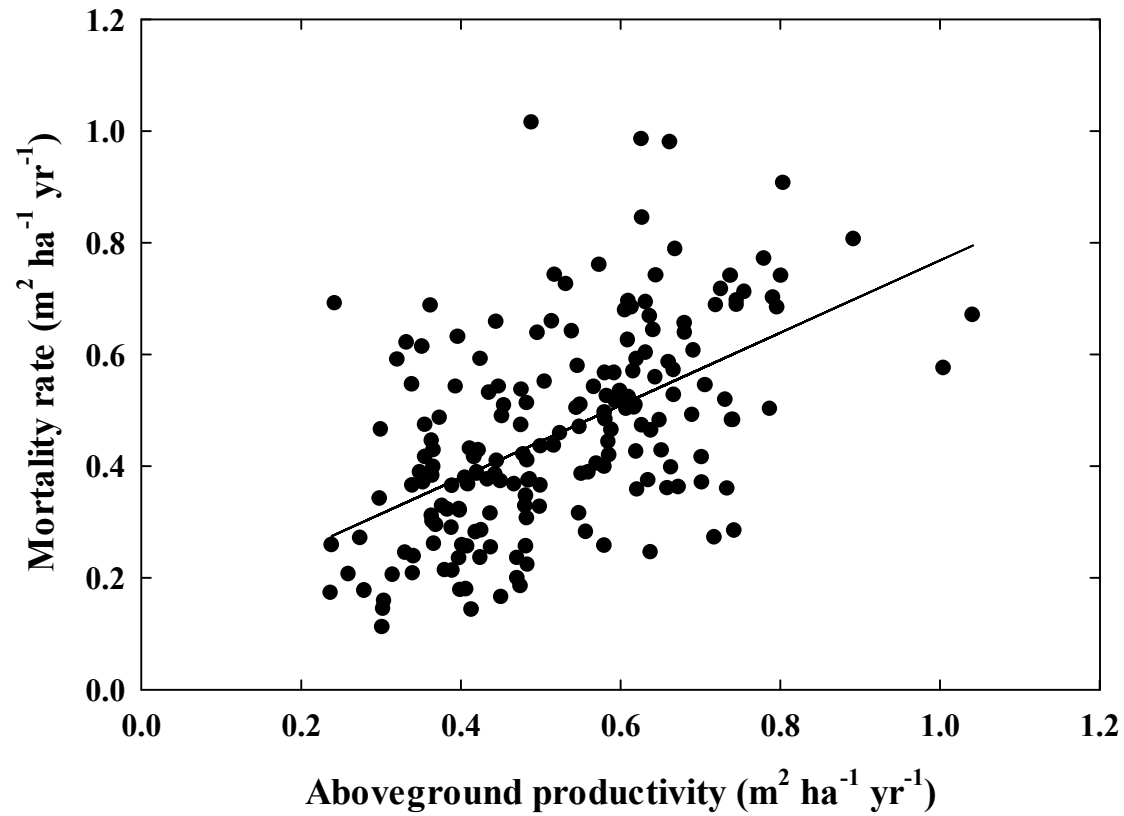


Figure 2. Consistent increases in mortality rate (% individuals died per total number of individuals per year) across the America's. Regression lines fitted to observations of stem mortality rate for the Amazon basin (solid line; Brien et al. 2015, slope of 0.029) and for temperate and boreal North America (dashed line; average values from all five sub-regions within van Mantgem et al. 2009, Peng et al. 2011, slope of 0.027). Linear regressions were used for simplicity, though a case can be made for non-linear (exponential) lines because zero-intercepts on the time-axis are not realistic (e.g. there is always some mortality occurring; see text). See supplemental information for methods details and for versions of this figure using different units.



737 **Figure 3: Basal area mortality rate is correlated with basal area productivity across the**
 738 **Amazon basin ($r^2 = 0.29$).** Data from Brien et al. 2015. Data represent stand dynamics as
 739 recorded for individual plots. See supplemental information for methods details.



740

Figure 4. A graphical summary of the literature evidence of changing mortality drivers and potential mechanisms over future conditions in moist -tropical forests. Shown are the expected trends in A) the forcing drivers of CO₂, temperature and VPD, and associated likelihoods of B) carbon starvation and/or hydraulic failure, C) liana abundance and fire frequency, D) biotic agent attack rates and destructive wind events, and E) competition for resources including shade. See references in text that support the general trends and their associated uncertainty. Panels C-E have widening uncertainty around the mean expectations due to lack of consistent projections (e.g. wind and biotic agents) or due to logical feedbacks (e.g. shade is enhanced by CO₂ but reduced by rising temperature and VPD; and CO₂ causes both

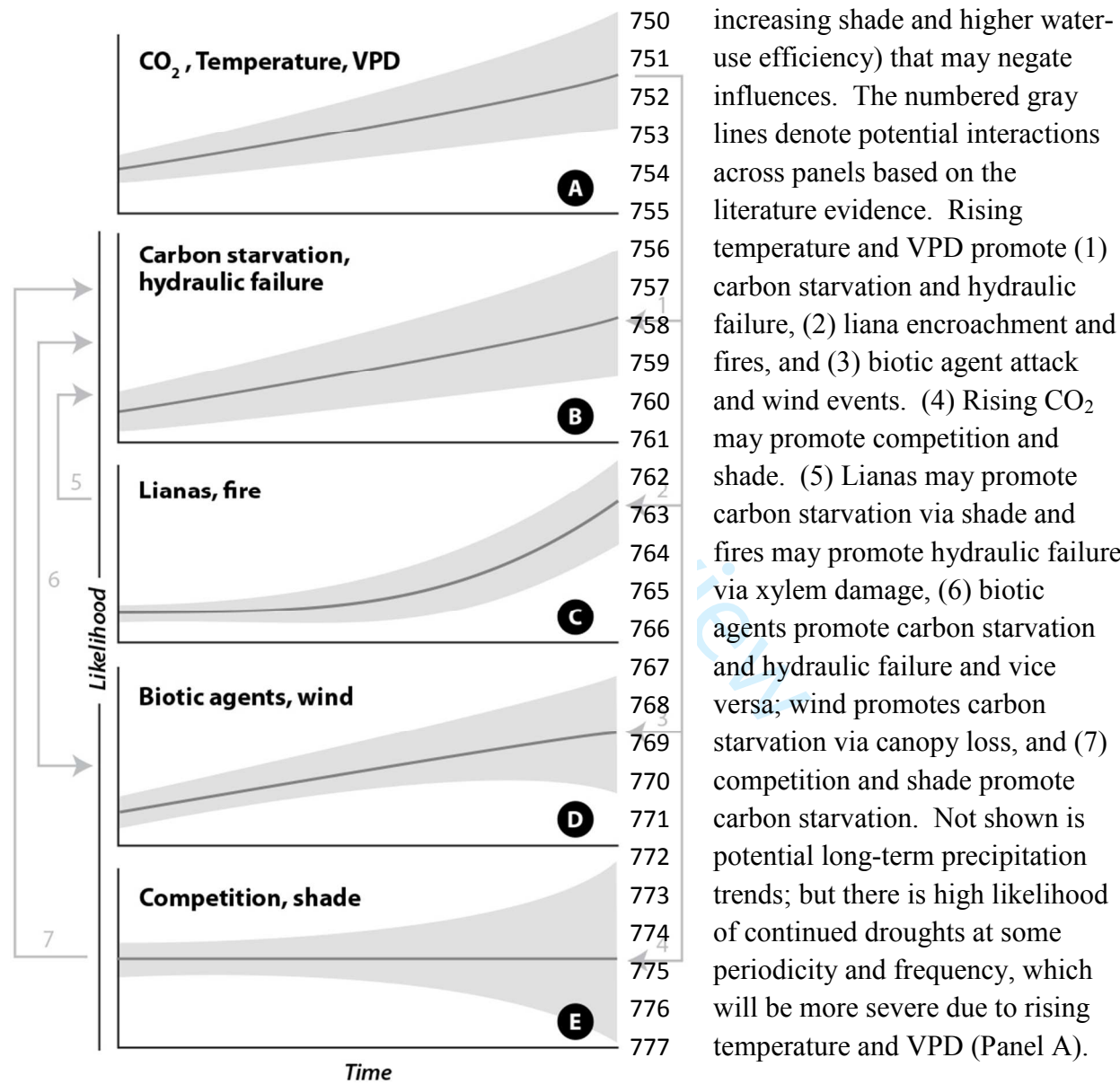
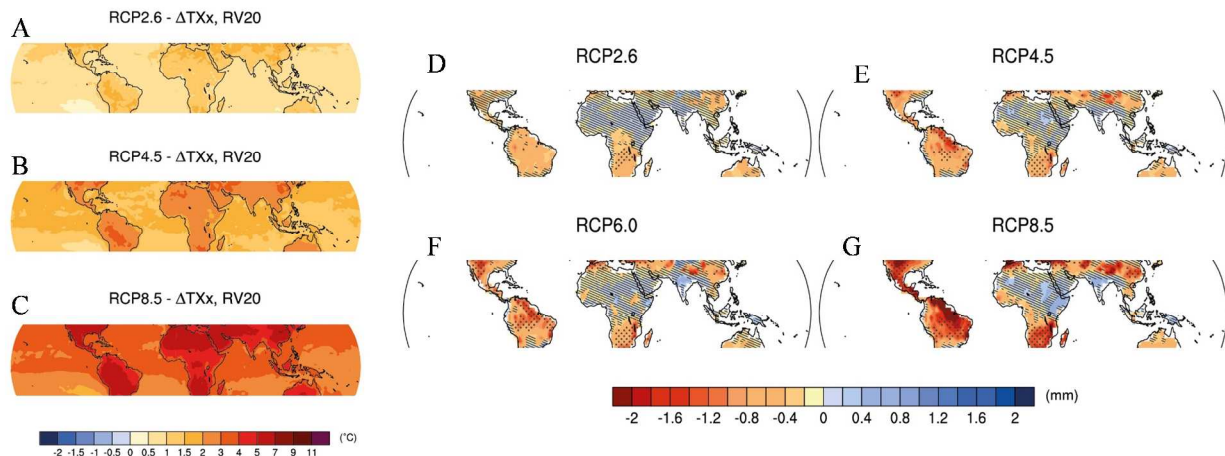


Figure 5. Coupled-Model Intercomparison Project (5) multi-model ensemble means of pan-tropical temperature and soil moisture in 2081-2100 relative to 1986-2005. **A-C)** The CMIP5 multi-model median change in 20-year return intervals of annual warm temperature extremes as simulated for 2081-2100 in RCP2.6 (top), RCP4.5 (middle), and RCP8.5 (bottom). **D-G)** Change in annual mean soil moisture (mass of water in the uppermost 10cm) (mm) for 2081-2100 relative to 1986-2005 from the CMIP5 ensemble (RCP2.6, 4.5, 6.0, and 8.5). Hatching indicates regions where the multi-model mean change is less than one standard deviation of internal variability and where at least 90% of models agree on the sign of change. Between 22 and 35 models were used depending on the scenario. Re-printed courtesy of Collins et al. (2013).



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Supplemental information

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- Figure SI3
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Figure 2 and Figure 3 Methods. Inventory data on the number of trees live and dead on a plot, censused at periodic intervals, can be converted into mortality rates using the equation

$$\text{mortality rate (year}^{-1}\text{)} = (d/n)/t \qquad \text{Eqn S1}$$

where n is the number of living trees at the start of the census interval, d is the number of those trees that died during the census interval, and t is the number of years in the census interval. However, this “arithmetic mortality” (as used by Brien et al. 2015) and also the commonly used “instantaneous mortality” rate (Eq S2), systematically underestimate mortality rate and can introduce systematic biases if the lengths of the census intervals change through time (Sheil et al. 1995).

This bias can be reduced by calculating the true annual mortality rate as

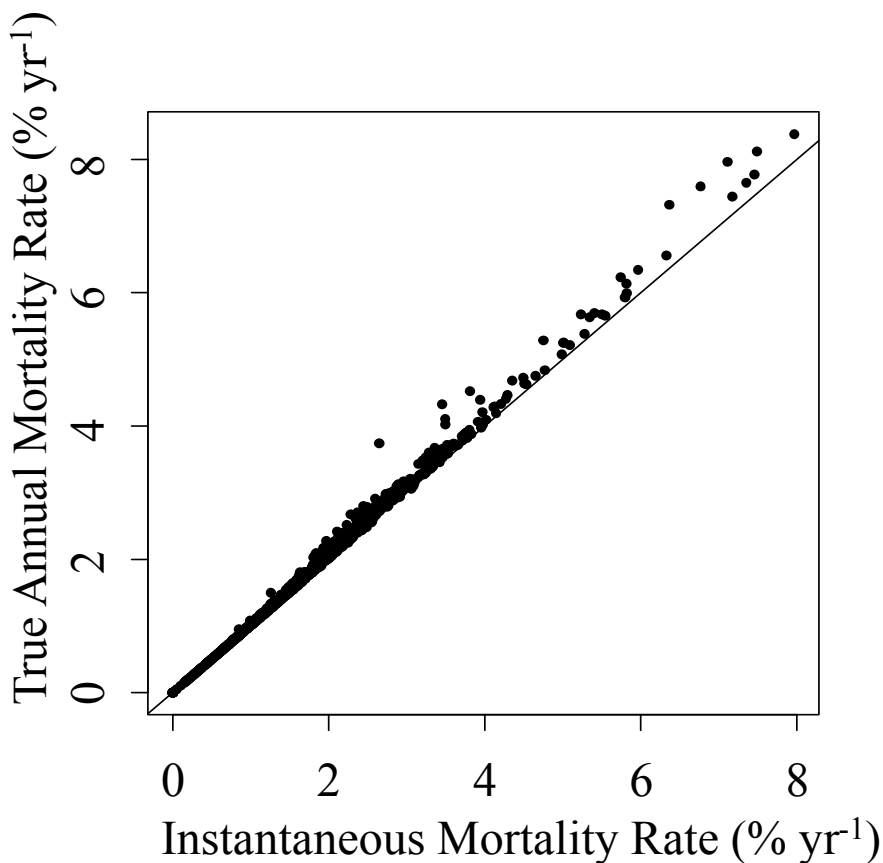
1646 mortality rate (year^{-1}) = $(\text{LN}(n) - (\text{LN}(n - d))/t$ Eq. S2.

1647

1648 (equation 4 in Sheil et al. 1995). The difference in estimates is generally small in absolute terms,

1649 especially when mortality rates are low (e.g., Figure SII).

1650



1651

1652 **Figure SII.** Comparison of the true annual mortality rate (Eq. S2) and the rate calculated under
 1653 the arithmetic formula (Eq. S1) for the Amazon tree dataset analyzed by Brien et al. (2015).
 1654 The mortality rate using Eq S1 is virtually indistinguishable from this result and is therefore not
 1655 shown. Each point represents a single plot and census interval. The line is the 1:1 line.

1656 Alternative versions of the Amazon data shown in Figure (2) are presented below in
1657 Figure SI2, including biomass mortality, percent mortality with all data shown, and percent
1658 biomass mortality (panels A, B, C respectively). Amazon mortality trends shown in Figure 2 are
1659 based on census data from 321 plots from across the Amazon basin from Brien et al. (2015).
1660 The trends are calculated using linear mixed effects models (lme4 package: Bates et al. 2013)
1661 with weightings applied based on sampling effort (plot size and monitoring length) as described
1662 in Brien et al. (2015). Mortality rates were calculated using equation S2 (Sheil et al. 1995;
1663 Kohyama et al. 2017).

1664 We did not conduct a formal test to compare the Amazonian and the N. American lines.
1665 We chose not to conduct this test based on limitations associated with the different time periods
1666 of the two datasets, and issues associated with new sites being added to each dataset throughout
1667 the life of the dataset. Formal comparison of global inventory datasets is a large but important
1668 future challenge.

1669 To assess whether the long-term trends in mortality rates for Amazon trees better fit a
1670 linear or non-linear model, we compared two mixed effects model fits. First, we performed a
1671 simple regression of the mid-point of each census interval against the mortality rate for that
1672 interval, using the lme4 package (Bates et al. 2013). This model accounted explicitly for plot
1673 effects (as random effect in the model) and for variation in sampling effort (plot size and census
1674 interval length) as outlined in Brien et al. 2015. We then rerun the same model, but adding a
1675 quadratic term to the explanatory variable (mid-point of census interval), in effect creating a non-
1676 linear fit. The two mixed effect models were then compared using ANOVA.

The implications of different equations to fit long-term mortality data (e.g. Figure 2) are substantial. Four possible interpretations of these data could be taken.

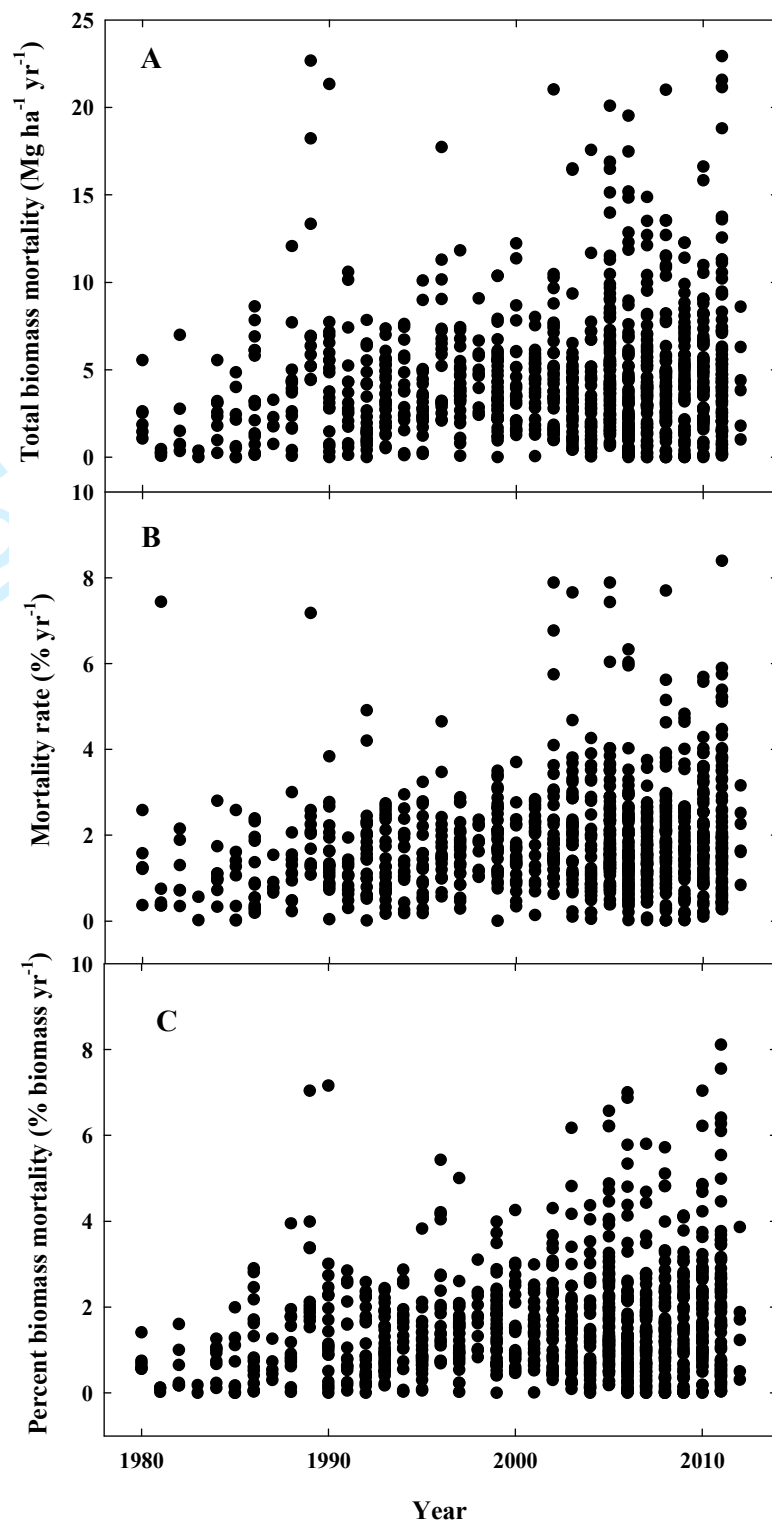
- 1) This time-window happened to capture an increase in mortality, but this is really part of long-term variation that is stationary. Mortality rates will continue to bounce around, with no fundamental change in their probabilities. .
- 2) The change in mortality rates during this time period is best fit by a step function; that is, mortality rates were previously stable at a lower rate, and transitioned or are transitioning to a new higher rates (say 2% instead of 1% for the Amazon). A doubling in mortality corresponds to a halving in residence time if all else is equal, thus this would have significant impacts on carbon storage.
- 3) Mortality rates increased linearly during this time period. If this increase continues for the foreseeable future, the rate will continue to rise as depicted in Figure (2), and we can expect another doubling in the near future. Clearly a linear increase cannot be extrapolated far backward in time, as mortality was never zero historically and cannot ever be negative.
- 4) Mortality rates increased exponentially during this time period. An exponential function is the best fit at low mortality rates (e.g. decades ago in Figure 2). Such an increase could be extrapolated backwards in time (it will never reach zero), although we lack data to support such extrapolation. Continued exponential growth in mortality would result in massive reductions in turnover time and biomass stocks.

Ultimately, understanding the proper relationships to fit to these long-term mortality data with an upward trend over time is more than simply a statistical exercise, but has implications for our understanding of the trajectory of mortality.

1700

For Peer Review

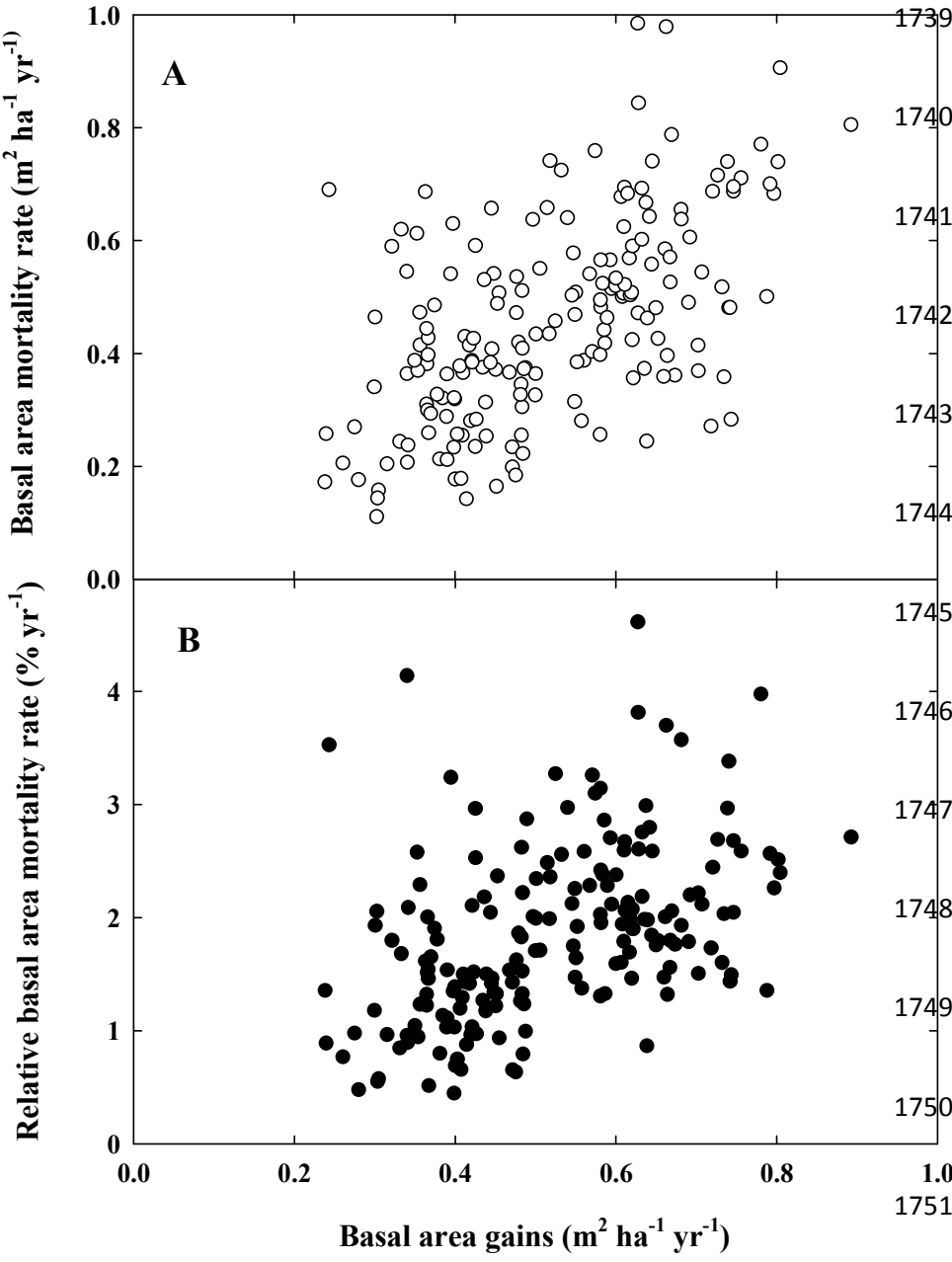
Figure SI2. Alternative approaches to presenting % mortality in units of # of individuals died / total number of individuals / year is to calculate the mortality in rates of biomass loss (Brienen et al. 2015). For completeness, we plotted multiple metrics of mortality using the Brienen et al. (2015) dataset in Figure SI1: A) biomass mortality (arithmetic equation SI1), B) % mortality, as shown in Figure (2) but with all data included to highlight both the trend and the variability, and C) percent of biomass lost to mortality. Census dates were rounded to the nearest year.



The relationship between mortality and productivity shown in Fig. 3 was based on the same set of Amazon plot data from Brien et al. (2015). Mortality rates of individual plots were calculated as the basal area of trees that died between censuses allowing for census interval effects (see Talbot et al. 2014). Plot level productivity was calculated as the sum of basal area growth of surviving trees, plus trees that recruited (that is reached > 100 mm in diameter) between censuses, and also included interval corrections as described in Talbot et al. 2014). We used a standard major axis regression (SMATR package: Warton and Weber 2002) to account for the errors in both productivity and mortality (x and y). The North American data are reproduced from van Mantgem et al. 2009 and Peng et al. 2011.

Alternative presentations of Figure (3) are shown in Figures SI3A, B.

Figure SI3. The relationship between mortality and productivity can be shown as it is in Figure (3) or in units of basal area gain (for productivity) and loss (for mortality), which more closely approximate biomass fluxes in absolute values (panel A) or relative to the total stand basal area (panel B).



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On nutrients

It is possible that elevated nitrogen availability is also driving increased growth and hence mortality. Measurements in Thailand and Panama have found that increased nitrogen deposition was associated with increased plant tissue nitrogen concentrations and nitrogen cycling rates without increasing productivity (Hietz et al. 2011). N deposition reduces cation availability and may thereby increase tree stress. N (and other particulate) deposition may increase or decrease the likelihood of mortality (Dietze and Moorcroft 2011, Gessler et al. 2017). No literature was available regarding phosphorous and mortality. Thus, atmospheric deposition may be partially responsible for trends in tree mortality, but its impact remains poorly studied in the tropics.

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1785 **A potential new approach to ESM modeling of hydraulics:** Inclusion of plant
1786 hydraulics in ESMs (or schemes capable of incorporation in ESMs) is drawing increased
1787 attention (Hickler et al. 2006, Alton et al. 2009, McDowell et al. 2013; Bonan et al. 2014,
1788 Mirfenderesgi et al., 2016, Christoffersen et al., 2016; Xu et al. 2016). These approaches apply
1789 Darcy's law to the coupled plant-soil continuum, which states that the rate of water supplied
1790 from the soil to the leaves is the product of the conductance in the xylem and the soil-leaf water
1791 potential gradient. A common theme to emerge from these model development studies is the
1792 improved realism of simulations, either with respect to transpiration at seasonal and interannual
1793 timescales, ecosystem response to water deficits, or drought-deciduous leaf phenology. A
1794 common challenge faced by any process-based representation of plant hydraulics in ESMs is
1795 maintaining a minimum level of complexity sufficient for capturing the first-order effect that
1796 plant hydraulics has on modulating ecosystem responses to water deficits. Sperry and Love
1797 (2015) have proposed a novel approach, attractive for ESMs, in which a simple integral
1798 transform is used to integrate the variable hydraulic properties from roots to stems and leaves to
1799 derive a representation of water delivery to the site of transpiration (the 'supply function'). A
1800 corresponding 'demand function' (and the ensuing modeled stomatal response to water deficits)
1801 then follows from theory suggesting that stomata operate in such a way to prevent catastrophic
1802 xylem failure, in which case hydraulic failure occurs through prolonged cuticular loss of water.
1803 This pragmatic approach to plant hydraulics well-simulates observed drought responses in
1804 tropical trees (Sperry and Love 2015). A recent extension of this work incorporates stomatal
1805 optimization of photosynthetic profit relative to hydraulic cost (Sperry et al. 2017), and
1806 provides an example for how hydraulics can potentially simplify simulating responses to a wide
1807 range of environmental cues (i.e., CO₂, light, temperature, as well as water deficits). In summary,

all evidence points to the inclusion of plant hydraulics in ESMs as a promising avenue for developing more mechanistic bases for tree mortality in the tropics.

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For Peer Review

Figure SI3. Projected changes in near-surface relative humidity percentage from CMIP5 models under RCP8.5 for the December, January, and February period (DJF, left), June, July, and August period (JJA, middle), and annual mean (ANN, right) averages relative to 1986-2005 for the periods 2046-2065 (top row) and 2081-2100 (bottom row). Hatching indicates regions where the multi-model mean change is less than one standard deviation of internal variability. Stippling indicates regions where the multi-model mean change is less than two standard deviations of internal variability and where 90% of models agree on the sign of change. Reprinted courtesy of Collins et al. (2013).

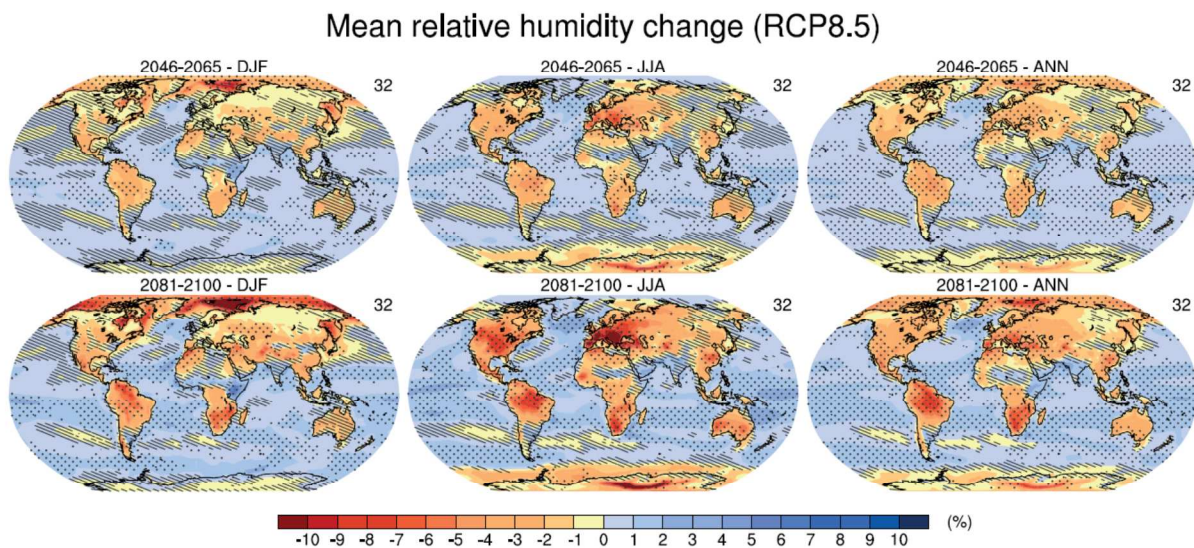


Figure SI4. CMIP5 multi-model ensemble average of zonal and annual mean wind change (2081-2100 minus 1986-2005) for, from left to right, RCP2.6, 4.5, and 8.5. Black contours represent the average for 1986-2005. Hatching indicates regions where the multi-model mean change is less than one standard deviation of internal variability, and stippling indicates regions where the multi-model mean change is greater than two standard deviation of internal variability and where at least 90% of models agree on the sign of change. Re-printed courtesy of Collins et al. (2013).

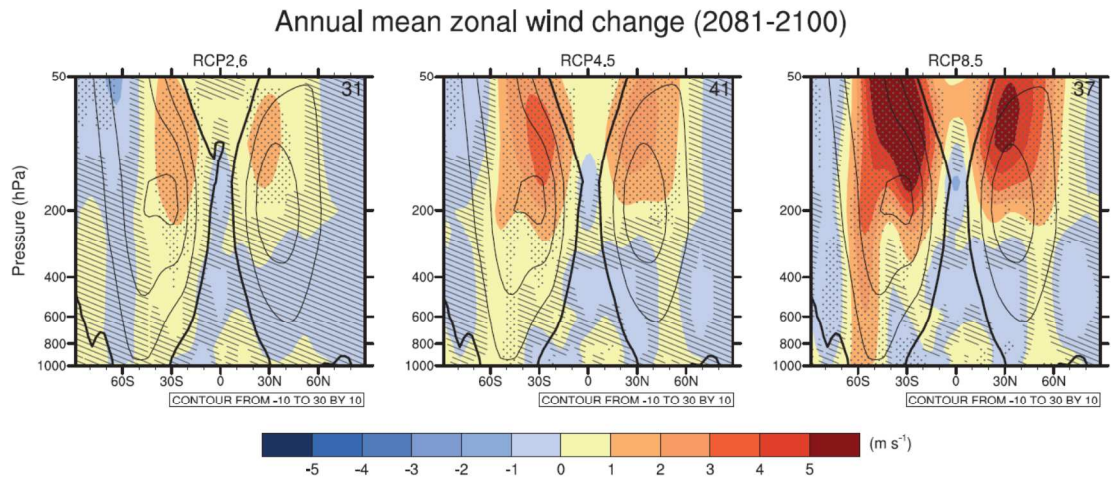


Figure SI5. The CMIP5 multi-model average percentage change in seasonal mean precipitation relative to the reference period 1986-2005 averaged for 2045-2065, 2081-2100, and 2181-2100 under the RCP8.5 forcing scenario. Hatching indicates regions where the multi-model mean change is less than one standard deviation of internal variability. Stippling indicates regions where the multi-model mean change is greater than two standard deviations of internal variability and where at least 90% of models agree on the sign of change. The number of models used in the analyses are shown in the upper right-hand corner of each figure. Re-printed courtesy of Collins et al. (2013).

